The limit that the true that the limit that the lim

The first term that the first term

	P									
EXPRESS MAIL LARFI	NI IMRED EI 610300881	US APR 20 O 2 APR 20 O S APR 20 O								
Form PTO-1390 U.S. DEPARTMENT OF COM	ATTORNEYS DOCKET NUMBER									
REV.10-94)	MENDE IN LETT MAD TO DE LA MANTE CONTROL	RN98131								
TRANSMITTAL LETTER TO TH	U.S. APPLICATION NO. (FINNWAL SEE 3737 CFR 15)									
DESIGNATED/elected office (De	NOT 10 19 19 18 10 16 16 19									
CONCERNING A FILING UNDE										
PCT/FR99/02345	October 1st, 1999	PRIORITY DATE CLAIMED October 2nd, 1998								
TITLE OF INVENTION										
DENTAL COMPOSITION BASED ON SILICONE CROSSLINKABLE BY										
CATION PROCESS										
APPLICANT (S) FOR DO/EO/US:										
Jean-Marc FRANCES										
Applicant herewith submits the United S	States Designated/Elected Office (De	D/EO/US) the following items and other								
information: 1 ☑ This is a FIRST submission of items concerning a filing under 35 U S C 371										
2 This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U S C 371										
3 🗵 This express request to begin national examination procedures (35 U S C 371) (f) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U S C 371 (b) and PCT Articles 22 and 39(1)										
4 A proper Demand for Internation										
claimed priority date										
5 🖂 A copy of the International Application as filed (35 U S C 371(c)(2))										
 a. ☐ is transmitted herewith (required only if not transmitted by the International Bureau). b. ☒ has been transmitted by the International Bureau. 										
c. is not required, as the application was filed in the United States Receiving Office (RO/US)										
6 A translation of the International Application into English (35 U S C 371(c)(2)).										
7 🛛 A copy of the International Search Report (PCT/ISA/210)										
8 🔯 Amendments to the claims of the International Application under PCT Article 19 (35 U S C 371(c)(2)). a. are transmitted herewith (required only if not transmitted by the International										
b. ☐ have been transmitted by the International Bureau										
c. ☐ have not been made, however the time limit for making such amendments has NOT expired. d. ☑ have not been made and will not be made.										
9 A translation of the amendments to the claims under PCT Article 19 (35 U S C 371(c)(3)).										
10 ⊠ A oath or declaration of the inventor(s) (35 U S C 371(c)(4)). (unsigned)										
11 🛛 A copy of the International Preliminary Examination Report (PCT/IPEA/409).										
12 A translation of the annexes to the International Preliminary Examination Report under PCT Article 36										
(35 U S C 371(c)(5)). Items 13 to 18 below concern document(s) or information included:										
13 An information Disclosure State	ment under 37 CFR 1 97 and 1 98									
14 An assignment document for rec	cording A separate cover sheet in c	ompliance with 37 CFR 3 28 and 3 31 is								
included 15 A FIRST preliminary amendmen	nt .									
16 A SECOND or SUBSEQUENT	F preliminary amendment									
17 A substitute specification.										
18 A change of power of attorney and/or address letter										
19 🔀 Certificate of Mailing by Express Mail										
20 Other items or information										
Forms PCT/IB/308 and 332, PC PCT/IPEA/ 409, PCT/FR99/023		ch Version),								

Page 1 of 2

C/Data/RN98131.Trans.doc

JC08 Rec'd PCT/PTO 0 2 APR 2001

US APPLICATION	976629		International Applic PCT/FR99/023		n No.		TORNEY DOCKET NUMBER N98131		
						CALCULATIONS PTO ONLY			
21 The following fees are submitted: BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)): Neither international preliminary examination fee (37 CFR 1,482) nor Intenational search fee (37 CFR 1.445(a)(2) paid to USPTO and International Search Report not prepared by the EPO or JPO \$1,000.00									
	ninary examination fee not paid th Report prepared by the EPC				860.00				
	inary examination fee (37 CFR arch (37 CFR 1,4445 (a)(2)) fe			\$	710.00				
☐ International preliminary examination fee paid to USPTO (37 CFR 1,482) but all claims did not satisfy provisions of PCT Article 33(1)-(4) \$ 690.00									
International preliminary examination fee paid to USPTO (37 CFR 1,482) and all claims satisfied provisions of PCT Article 33(2)-(4)									
Surcharge of \$130.00 for furnishing the oath or declaration later than 20 30 months from the earliest claimed priority date (37 CFR 1,492(e)).						\$	0.00		
GLAIMS	NUMBER FILED	NUN	IBER EXTRA	T RA	ATE .	*	0.00		
Total Claims	8 -20=		0		18.00	\$	0.00	[
Independent Claims	1 -3=		0	X\$	80.00	\$	0.00		
	NT CLAIMS(S) (if applicable)			1 []	\$	0.00		
TOTAL OF ABOVE CALCULATIONS = Reduction by 1/2 for filing by small entity, if applicable. Verified Small Entity Statement must also						\$	0.00	 	
be filed (Note 37 CFR 1.9, 1.27, 1.28)						L.			
SUBTOTAL = Processing fee of \$130.00 for furnishing the English translation later than 20 30 months						\$	860.00		
from the earliest claimed priority date (37 CFR 1,492 (f)).						\$	0.00		
TOTAL NATIONAL FEE = Fee for recording the enclosed assignment (37 CFR1.21(h)). The assignment must be						\$	860.00	 	
accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property + 🗵 \$ 0.00									
TOTAL FEES ENCLOSED =						\$	860.00		
\$ 000 \$ 2 0 2 000 2 000						Amount to be			
						refunded \$ charged \$			
ner						cna	rgea 	*	
a A check in the amount of \$ to cover the above fees is enclosed									
b 🔯 Please charge my Deposit Account No. 18-1171 in the amount of \$860.00 to cover the above fees.									
c The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 18-1171 A duplicate copy of this sheet is enclosed.									
NOTE: Where an appropriate time limit under 37 CFR 1,494 or 1,495 has not been met, a petition to revive (37 CFR 1.137 (a) or (b)) must be filed and granted to restore the application to pending status.									
SEND ALL CORRESPONDENCE TO: Jean-Louis SEUGNET SIGNATURE									
RHODIA INC. Jean-Louis SEUGNET									
259 Prospect Plains Road NAME CN 7500 Limited Recognition under 37 C						ED S	10 9/h)		
Cranbury, NJ 08512	Cranbury, NJ 08512 enclosed.								
April 02, 2001 REGISTRATION NUMBER									

JC08 Rec'd PCT/PTO

Case RN98131

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In the Application of: Jean-Marc FRANCES

National Phase of PCT/FR99/02345

Examiner: N/A

International Filing Date: October 1st, 1999

Art Unit: N/A

Serial No: N/A

Filing Date: April 2, 2001

For: DENTAL COMPOSITION BASED ON SILICONE CROSSLINKABLE

BY CATION PROCESS

Assistant Commissioner for Patents Washington, D.C. 20231

PRELIMINARY AMENDMENT

Dear Sir:

Prior to calculation of filing fee, please enter the following amendment in the specification and claims:

In the specification:

Page 1, just after the title, please add: - - This application is an application under 35 U.S.C. Section 371 of International Application Number PCT/FR99/02345 filed on

October 1st, 1999.- -

In the claims:

Please cancel claims 1 to 11 and replace them with the following new claims 12 to 19:

- 12. (New) A dental composition comprising:
- (1) at least one crosslinkable or polymerizable silicone oligomer or polymer which is liquid at room temperature or which is heat-meltable at a temperature of less than 100°C, and which comprises:

a) at least one unit of formula (FS):

$$Z - Si - R^0 \rightarrow O_{(3-a)/2}$$

wherein:

$$a = 0, 1 \text{ or } 2,$$

- R⁰, identical or different, represents an alkyl, cycloalkyl, aryl, vinyl, hydrogeno or alkoxy radical,
- Z, identical or different, is an organic substituent comprising at least one reactive epoxy, alkenyl ether, oxetane, dioxolane or carbonate functional group, and
 - b) at least two silicon atoms,
- (2) at least one aromatic hydrocarbon photosensitizer with one or more aromatic nuclei which are optionally substituted, and having a residual light absorption of between 200 and 500 nm,
- (3) at least one dental filler present in a proportion of at least 10% by weight relative to the total weight of the composition, and
 - (4) an effective quantity of at least one borate photoinitiator having an anionic borate entity and a cationic borate entity and whose cationic entity is:

onium salts of formula (I):

$$[(R^1)_n - A - (R^2)_m]^+ (I)$$

in which formula:

A represents an element of groups 15 to 17 of valency v,

 R^1 represents a carbocyclic or heterocyclic C_6 - C_{20} aryl radical, optionally containing a nitrogen or sulfur atom,

 R^2 represents R^1 or a linear or branched C_1 - C_{30} alkyl or alkenyl radical; said radicals R^1 and R^2 being optionally substituted with a C_1 - C_{25} alkoxy, C_1 - C_{25} alkyl, nitro, chloro, bromo, cyano, carboxyl, ester or mercapto group, n is an integer ranging from 1 to v+1, v being the valency of the element A, m is an integer ranging from 0 to v-1 with n+m=v+1,

(**) organometallic salts of formula (III):

$$(L^1L^2L^3M)^{+q}$$

in which formula:

M represents a group 4 to 10 metal,

 L^1 represents 1 ligand bound to the metal M by π by electrons, said ligand being η^3 -alkyl, η^5 -cyclopendadienyl, η^7 -cycloheptratrienyl, optionally substituted η^6 -aromatic compounds, or compounds having from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 3 to 8 π electrons;

 L^2 represents a ligand bound to the metal M by π electrons, said ligand being η^7 -cycloheptratrienyl, optionally substituted η^6 -benzene, or compounds having

from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 6 or 7 π electrons;

 L^3 represents from 0 to 3 ligands, which are identical or different, linked to the metal M by σ electrons, saidligand(s) being CO, or NO_2^+ ; the total electron charge q of the complex to which L^1 , L^2 and L^3 contribute and the ionic charge of the metal M being positive and equal to 1 or 2; and

whose anionic borate entity is of the formula [BX_aR_b] wherein:

- a and b are integers ranging, for a, from 0 to 3 and, for b, from 1 to 4 with a + b = 4,
- the symbols X represent:
 - a halogen atom (chlorine, fluorine) with a = 0 to 3,
 - an OH functional group with a = 0 to 2,
- the symbols R are identical or different and represent:
 - a phenyl radical substituted with at least one electron-attracting group or with at least 2 halogen atoms, when the cationic entity is an onium of an element of groups 15 to 17,
 - a phenyl radical substituted with at least one element or one electron-attracting group, when the cationic entity is an organometallic complex of an element of groups 4 to 10,

an aryl radical containing at least two aromatic nuclei optionally substituted with at least one electron-attracting group or element, regardless of the cationic entity.

- 13. (New) A composition as claimed in claim 12, wherein Z is an organic substituent Z1 comprising at least one reactive epoxy, or dioxolane functional group.
- 14. (New) A composition as claimed in claim 13, wherein the oligomer or polymer (1) further comprises other reactive functional groups Z which are alkenyl ether, oxetane or carbonate functional groups Z2.
- 15. (New) A composition as claimed in claim 12, wherein the reactive functional group(s) of Z1 are selected from the group consisting of the following radicals:

16. (New) A composition as claimed in claim 12, wherein the photoinitiator is

17. (New) A composition as claimed in claim 12, wherein the photosensitizer

is:

4,4'-dimethoxybenzoin; 2-4-diethylthioxanthone

2-ethylanthraquinone; 2-methylanthraquinone;

1,8-dihydroxyanthraquinone; dibenzoylperoxide;

2,2-dimethoxy-2-phenylacetophenone;

benzoin;

2-hydroxy-2-methylpropiophenone;

benzaldehyde;

4-(2-hydroxyethoxy)phenyl-(2-hydroxy-2-methylpropyl)-ketone;

benzoylacetone;

2-isopropylthioxanthone;

1-chloro-4-propoxythioxanthone; or

4-isopropylthioxanthone.

18. (New) A dental composition as claimed in claim 12, wherein the silicone oligomer or polymer (1) comprises at least one silicone having the following average formula:

or

CHCH CH2

CH2CH2CH2

CH3 Si-CH3

CH3 Si-CH3

CH2CH2CH2

CH2CH2CH2

19. (New) A process for the preparation of a dental prosthese or dental restoration, comprising the step of using a dental composition as defined in claim 12.

REMARKS

It is asserted that these amendments do not add new matter. Support for these amendments can be found in the specification and claims as originally filed. Entry of these amendments is respectfully requested.

April 67, 2001

A STATE OF THE STA

111 122

Min and

i D

RHODIA INC. 259 Prospect Plains Road CN7500, Cranbury, NJ 08512

F/Brevet/RN98131..prelim

Respectfully submitted

JEAN-LOUIS SEUGNET

Limited Recognition under 37 CFR § 10.9(b)

enclosed

Tel: (609) 860-4180 Fax: (609) 860-0503

SULBBPPEPLAJ3

- 1 -

PCT/FR99/02345

DENTAL COMPOSITION BASED ON SILICONE CROSSLINKABLE BY CATION PROCESS

The field of the invention is that of dental compositions. More precisely, the dental compositions

5 used in the context of the present invention can be used for producing dental prostheses and for dental restoration.

Up until now, to produce dental compositions for the preparation of dental prostheses or of dental restoration materials, it is possible to use resins based on photopolymerizable acrylates. These ready-to-formulate products exhibit however upon use problems of irritation and potential problems of toxicity.

In addition, these products have the major

disadvantage of causing high volume shrinkage during
their polymerization, which makes their use complex and
difficult for the production of dental prostheses or of
dental restoration materials. Problems of attachment
due to the volume shrinkage or to the lack of adherence
of the polymers used are in particular observed.

The object of the present invention is to provide novel dental compositions which do not exhibit the disadvantages of the prior art. These novel dental compositions, which are polymerizable and/or

crosslinkable in an oral environment, have markedly improved qualities, in particular as regards the very marked reduction in the phenomenon of shrinkage of the

dental compositions used for the production of dental prostheses or of dental restoration materials.

The polymerizable and/or crosslinkable dental composition according to the invention comprises:

(1) at least one crosslinkable and/or polymerizable silicone oligomer or polymer which is liquid at room temperature or which is heat-meltable at a temperature of less than 100°C, and which comprises:

• at least one unit of formula (FS): $Z - Si - (R^0)_a O_{(3-a)/2}$

in which:

- -a = 0, 1 or 2,
- R^0 , identical or different, represents an alkyl, cycloalkyl, aryl, vinyl, hydrogeno or alkoxy radical, preferably a C_1 - C_6 lower alkyl,
- Z, identical or different, is an organic substituent comprising at least one reactive epoxy, and/or alkenyl ether and/or oxetane and/or dioxolane and/or carbonate functional group, and preferably Z being an organic substituent comprising at least one reactive epoxy and/or dioxolane functional group,
- and at least two silicon atoms,

10

5

15

20

25

10

- (2) an effective quantity of at least one borate-type photoinitiator,
- (3) at least one aromatic hydrocarbon photosensitizer with one or more aromatic nuclei which are substituted or not, having a residual light absorption of between 200 and 500 nm.
- (4) and at least one dental filler present in a proportion of at least 10% by weight relative to the total weight of the composition.

According to a first advantageous variant of the present invention, the dental composition is polymerizable and/or crosslinkable under activation by the thermal route and/or by the photochemical route.

In general, the photochemical activation is carried out under UV radiation. More particularly, UV radiation having a wavelength of the order of 200 to 500 nm is used for the production of dental prostheses and UV-visible radiation having a wavelength greater than 400 nm for the production of restoration materials. A wavelength greater than 400 nm allows crosslinking and/or polymerization in an oral environment.

25 The silicone polymer or oligomer (1) has the advantage, compared with organic resins which are crosslinked by the cationic route, of being transparent

to UV-visible light and therefore its use makes it possible to obtain materials which are very thick and whose photocrosslinking occurs in a short time.

The reactive functional groups Z of the

5 silicone polymer or oligomer (1) may be highly varied.

However, particularly advantageous dental compositions

are obtained when the silicone oligomer or polymer (1)

comprises at least one (FS) unit in which Z represents

an organic substituent Z1 comprising at least one

10 reactive epoxy, and/or dioxolane functional group, and

preferably at least one reactive epoxy functional

group.

According to two advantageous alternatives of the present invention, the silicone oligomer or polymer (1) with at least one reactive epoxy and/or dioxolane functional group Z1, and preferably at least one reactive epoxy functional group may:

- (i) either comprise only this (these) type(s)
 of reactive functional group(s) Z1,
- 20 (ii) or comprise other reactive functional groups Z such as the reactive alkenyl ether, oxetane and/or carbonate functional groups Z2.

In the case of the first alternative (i), the

25 dental composition may also comprise other silicone

oligomers and/or polymers comprising other reactive

functional groups Z2 such as alkenyl ether, oxetane

and/or carbonate functional groups and optionally reactive functional groups Z1.

By way of examples of reactive functional groups Z, these may in particular be chosen from the following radicals:

- with R" representing a linear or branched $C_1\text{-}C_6$ alkyl radical.

According to a second advantageous variant of the present invention, the silicone polymer or oligomer consists of at least one silicone having the following average formula:

k)
$$CH_3 \qquad CH_3 \qquad CH_3$$

$$CH_3 \qquad CH_3 \qquad CH_3$$

$$A < 1000.$$

1) $\frac{R_0}{a} = \frac{R_0}{a} = \frac{R_0}{a}$ a < 1000.

The cationic photoinitiators are chosen from onium borates (taken on their own or as a mixture with each other) of an element of groups 15 to 17 of the

5 Periodic Table [Chem. & Eng. News, vol. 63, No. 5, 26 of 4 February 1985] or of an organometallic complex of an element of groups 4 to 10 of the Periodic Table [same reference].

The cationic entity of the borate is selected 10 from:

(1) the onium salts of formula (I)

$$[(R^1)_n - A - (R^2)_m]^+$$
 (I)

in which formula:

- A represents an element of groups 15 to 17 such as for example: I, S, Se, P or N,
- R^1 represents a carbocyclic or heterocyclic C_6 - C_{20} aryl radical, it being possible for said heterocyclic radical to contain, as heteroelements, nitrogen or sulfur,

15

- R^2 represents R^1 or a linear or branched C_1 - C_{30} alkyl or alkenyl radical; said radicals R^1 and R^2 being optionally substituted with a C_1 - C_{25} alkoxy, C_1 - C_{25} alkyl, nitro, chloro, bromo, cyano, carboxyl, ester or mercapto group,
- n is an integer ranging from 1 to v + 1, v being the valency of the element A,
- m is an integer ranging from 0 to v-1 with n+m=v+1,
- (2) the oxoisothiochromanium salts described in patent application WO 90/11303, in particular the sulfonium salt of 2-ethyl-4-oxoisothiochromanium or 2-dodecyl-4-oxoisothiochromanium,
 - (3) the sulfonium salts in which the cationic entity comprises:
 - 3₁ at least one polysulfonium entity of formula (II.1):

$$Ar^{1} - S - Ar^{3} - Y - \begin{bmatrix} Ar^{3} - \overline{S} - Ar^{1} \\ Ar^{2} \end{bmatrix}$$

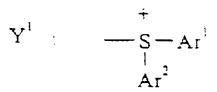
in which:

20 - the symbols Ar^1 , which are identical or different, each represent a monovalent phenyl or naphthyl radical, optionally substituted with one or more radicals chosen from: a linear or branched C_1-C_{12} , preferably C_1-C_6 , alkyl radical, a linear or branched C_1-C_{12} , preferably

 C_1 - C_6 , alkoxy radical, a halogen atom, an -OH group, a -COOH group, an ester group -COO-alkyl where the alkyl portion is a linear or branched C_1 - C_{12} , preferably C_1 - C_6 , residue, and a group of formula -Y⁴-Ar² where the

- 5 symbols Y^4 and Ar^2 have the meanings given just below,
 - the symbols Ar^2 , which are identical or different, each represent a monovalent phenyl or naphthyl radical, optionally substituted with one or more radicals chosen from: a linear or branched C_1 - C_{12} , preferably C_1 - C_6 ,
- alkyl radical, a linear or branched C_1-C_{12} , preferably C_1-C_6 , alkoxy radical, a halogen atom, an -OH group, a -COOH group, an ester group -COO-alkyl where the alkyl portion is a linear or branched C_1-C_{12} , preferably C_1-C_6 , residue,
- 15 the symbols Ar^3 , which are identical or different, each represent a divalent phenylene or naphthylene radical, optionally substituted with one or more radicals chosen from: a linear or branched C_1 - C_{12} , preferably C_1 - C_6 , alkyl radical, a linear or branched
- 20 C_1 - C_{12} , preferably C_1 - C_6 , alkoxy radical, a halogen atom, an -OH group, a -COOH group, an ester group -COO-alkyl where the alkyl portion is a linear or branched C_1 - C_{12} , preferably C_1 - C_6 , residue,
- t is an integer equal to 0 or 1, with the additional conditions according to which:
 - + when t = 0, the symbol Y is then a monovalent radical Y^1 representing the group of formula:





where the symbols Ar^1 and Ar^2 possess the meanings given above,

5 + when t = 1:

 $oldsymbol{\circ}$ on the one hand, the symbol Y is then a divalent radical having the following meanings Y² to Y⁴:

• Y²: a group of formula:



where the symbol Ar² has the meanings given above,

• Y3: a single valency bond,

• Y4: a divalent residue chosen from:

a linear or branched C_1-C_{12} , preferably C_1-C_6 , alkylene residue and a residue of formula $-\text{Si}(CH_3)_2\text{O-}$,

oldsymbol on the other hand, in the case solely where the symbol Y represents Y³ or Y⁴, the (terminal) radicals Ar¹ and Ar² possess, in addition to the meanings given above, the possibility of being linked to each other by the residue Y' consisting of Y'¹ a single valency bond or of Y'² a divalent

residue chosen from the residues cited in relation to the definition of Y^4 , which is inserted between the carbon atoms, opposite each other, situated on each aromatic ring at the ortho position with respect to the carbon atom directly linked to the cation S^+ ;

• 3_2 and/or at least one monosulfonium entity possessing a single cationic center S^+ per mol of cation and consisting, in most cases, of entity of formula (II.2):

$$Ar^{1} - S - Ar^{3}$$

$$Ar^{2}$$

10

5

in which Ar^1 and Ar^2 have the meanings given above in relation to formula (III.1), including the possibility of directly linking to each other only one of the radicals Ar^1 to Ar^2 in the manner indicated above in relation to the definition of the additional condition in force when t=1 in formula (II), calling into play the residue Y';

(4) the organometallic salts of formula (III):

$$(L^1L^2L^3M)^{+q}$$

20 in which formula:

- M represents a group 4 to 10 metal, in particular iron, manganese, chromium or cobalt,
- L¹ represents 1 ligand bound to the metal M by π electrons, which ligand is chosen from the ligands η^3 -alkyl, η^5 -cyclopendadienyl and η^7 -cycloheptratrienyl

and the η^6 -aromatic compounds chosen from the optionally substituted η^6 -benzene ligands and the compounds having from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 3 to 8 π electrons;

- L^2 represents a ligand bound to the metal M by π electrons, which ligand is chosen from the ligands η^7 -cycloheptratrienyl and the η^6 -aromatic compounds chosen from the optionally substituted ligands 10 η^6 -benzene and the compounds having from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 6 or 7 π electrons;
- L^3 represents from 0 to 3 ligands, which are identical or different, linked to the metal M by σ electrons, which ligand(s) is (are) chosen from CO and NO_2 , the total electron charge q of the complex to which L^1 , L^2 and L^3 contribute and the ionic charge of the metal M being positive and equal to 1 or 2;
- The anionic borate entity has the formula $[BX_aR_b]^-$ in which:
 - a and b are integers ranging, for a, from 0 to 3 and, for b, from 1 to 4 with a + b = 4,
 - the symbols X represent:
- * a halogen atom (chlorine, fluorine) with a = 0
 to 3,
 - * an OH functional group with a = 0 to 2,

10

- the symbols R are identical or different and represent:

▶ a phenyl radical substituted with at least one electron-attracting group such as for example OCF₃, CF₃, NO₂, CN, and/or with at least 2 halogen atoms (fluorine most particularly), this being when the cationic entity is an onium of an element of groups 15 to 17,

➤ a phenyl radical substituted with at least one element or one electron-attracting group, in particular a halogen atom (fluorine most particularly), CF₃, OCF₃, NO₂, CN, this being when the cationic entity is an organometallic complex of an element of groups 4 to 10,

an aryl radical containing at least two aromatic nuclei such as for example biphenyl, naphthyl, optionally substituted with at least one electron-attracting group or element, in particular a halogen atom (fluorine most particularly), OCF3, CF3, NO2, CN, regardless of the cationic entity.

Without being limiting, more details are given below as regards the subclasses of onium borate and of borate of organometallic salts more particularly preferred in the context of the use in accordance with the invention.

According to a first preferred variant of the invention, the entity of the anionic borate entity which are most particularly suitable are the following:

 $5': [B(C_6H_3(CF_3)_2)_4]^-$ 1': $[B(C_6F_5)_4]^-$

2': $[(C_6F_5)_2BF_2]^-$ 6': $[B(C_6H_3F_2)_4]^-$

 $7': [C_6F_5BF_3]^ 3': [B(C_6H_4CF_3)_4]^-$

4': $[B(C_6F_4OCF_3)_4]^{-}$.

According to a second preferred variant of the invention, the onium salts (1) which can be used are described in numerous documents, in particular in patents US-A-4 026 705, US-A-4 032 673, US-A-4 069 056, US-A-4 173 476. US-A-4 136 102, Among these, 10 following cations will be most particularly preferred:

According to a third preferred variant, the organometallic salts (4) which can be used US-A-4 973 722, documents the 15 described in US-A-4 992 572, EP-A-203 829, EP-A-323 584 EP-A-354 181. The organometallic salts most readily selected according to the invention are in particular:

 $(\eta^5$ -cyclopentadienyl) $(\eta^6$ -toluene) Fe⁺,

 $(\eta^5$ -cyclopentadienyl) $(\eta^6$ -methyl-1-naphthalene) 20 Fe⁺,

```
.(\eta^5-cyclopentadienyl)(\eta^6-cumene) Fe<sup>+</sup>, .bis(\eta^6-mesitylene) Fe<sup>+</sup>, .bis(\eta^6-benzene) Cr<sup>+</sup>.
```

In agreement with these three preferred 5 variants, the following products may be mentioned by way of examples of photoinitiators of the onium borate type:

 $.\;(\eta^{5}\text{-cyclopentadienyl})\;(\eta^{6}\text{-toluene})\;\;\text{Fe}^{\scriptscriptstyle +},\;\;[\,\text{B}\,(\text{C}_{6}F_{5})_{\,4}\,]^{\scriptscriptstyle -}$

.(η^5 -cyclopentadienyl)(η^6 -methyl-1-naphthalene)

 Fe^+ , $[B(C_6F_5)_4]^-$

.(η^5 -cyclopentadienyl)(η^6 -cumene) Fe $^-$, [B(C $_6$ F $_5$) $_4$]

As another literary reference for defining the onium borates (1) and (2) and the borates of organometallic salts (4), there may be mentioned the entire content of patent applications EP 0 562 897 and 0 562 922. This content is integrally incorporated by reference into the present disclosure.

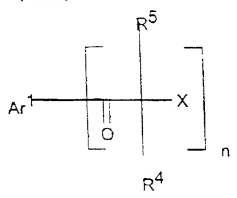
As another example of onium salt which can be used as photoinitiator, there may be mentioned those disclosed in American patents US 4 138 255 and US 4 310 469.

10

Other cationic photoinitiators may also be used, e.g.:

- those marketed by Union-Carbide (photoinitiator 6990 and 6974 triarylsulfonium hexafluorophosphate and hexafluoroantimonate),
- the salts of iodonium hexafluorophosphate or hexafluoroantimonate,
- or the ferrocenium salts of these various anions.

The nature of the photosensitizer contained in the dental composition according to the invention may be highly varied. In the context of the invention, it corresponds in particular to one of the following formulae (IV) to (XXII):



• formula (IV)

in which:

20 - when n = 1, Ar^1 represents an aryl radical containing from 6 to 18 carbon atoms, a

tetrahydronaphthyl, thienyl, pyridyl or furyl radical or a phenyl radical carrying one or more substituents chosen from the group consisting of F, Cl, Br, CN, OH, linear or branched C_1-C_{12} alkyls, $-CF^3$, $-OR^6$, -OPhenyl, $-SR^6$, -SPhenyl, $-SO_2$ Phenyl, $-COOR^6$, $-O-(CH_2-CH=CH_2)$, $-O(CH_2H_4-O)_m-H$, $-0(C_3H_6O)_m$ -H, m being between 1 and 100,

- when n = 2, Ar_1 represents a C_6-C_{12} arylene radical or a phenylene-T-phenylene radical where T represents -O-, -S-, -SO₂- or -CH₂-,
- \mathbf{X} represents a group $-\mathrm{OR}^7$ or $-\mathrm{OSiR}^8(\mathrm{R}^9)_2$ or forms, with R^4 , a group $-O-CH(R^{10})-$.
- R4 represents a linear or branched C1-C8 alkyl radical which is unsubstituted or which carries an -OH, $-OR^6$, C_2-C_8 acyloxy, $-CF^3$ or -CN group, a C_3 or C_4 alkenyl radical, a C_6 to C_{18} aryl radical, a C₇ to C₉ phenylalkyl radical,
- \mathbf{R}^{5} has one of the meanings given for \mathbf{R}^{4} or represents a radical -CH₂CH₂R¹¹, or alternatively forms with R^4 a C_2 - C_8 alkylene radical or a C₃-C₉ oxa-alkylene or aza-alkylene radical,
- R⁶ represents a lower alkyl radical containing from 1 to 12 carbon atoms,
- \mathbf{R}^7 represents a hydrogen atom, a C_1 - C_{12} alkyl radical, a C2-C6 alkyl radical carrying an -OH, $-OR^6$ or -CN group, a C_3-C_6 alkenyl radical, a cyclohexyl or benzyl radical, a phenyl radical

10

5

15

25

20

10

15

optionally substituted with a chlorine atom or a linear or branched $C_1 - C_{12}$ alkyl radical, or a 2-tetrahydropyranyl radical,

- $\mathbf{R^8}$ and $\mathbf{R^9}$ are identical or different and each represent a $C_1\text{-}C_4$ alkyl radical or a phenyl radical.
- \mathbf{R}^{10} represents a hydrogen atom, a C_1 - C_8 alkyl radical or a phenyl radical,
- \mathbf{R}^{11} represents a radical -CONH₂, -CONHR⁶, -CON(\mathbf{R}^{6})₂, -P(O)(OR⁶)₂ or 2-pyridyl;

Ar² R¹⁵

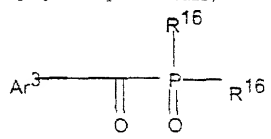
• formula (V)

in which:

- Ar^2 has the same meaning as Ar^1 of formula (IV) in the case where n=1,
- R^{15} represents a radical chosen from the group consisting of a radical Ar^2 , a linear or branched C_1 - C_{12} alkyl radical, a C_6 - C_{12} cycloalkyl radical, and a cycloalkyl radical forming a C_6 - C_{12} ring with the carbon of the ketone or a carbon of the radical Ar^2 , it being possible for these radicals to be substituted with one or more substituents chosen from the group consisting of -F, -Cl, -Br, -CN, -OH, -CF₃, -OR⁶, -SR⁶, -COOR⁶, the linear or branched C_1 - C_{12} alkyl radicals optionally carrying

15

an -OH, $-OR^6$ and/or -CN group, and the linear or branched C_1-C_8 alkenyl radicals;

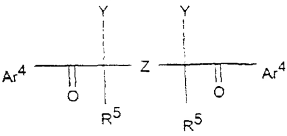


• formula (VI)

5 in which:

 \mathbf{Ar}^3 has the same meaning as \mathbf{Ar}^1 of formula (IV) in the case where $\mathbf{n}=1$,

- R^{16} , identical or different, represents a radical chosen from the group consisting of a radical Ar^3 , a radical $-(C=0)-Ar^3$, a linear or branched C_1-C_{12} alkyl radical, a C_6-C_{12} cycloalkyl radical, it being possible for these radicals to be substituted with one or more substituents chosen from the group consisting of -F, -Cl, -Br, -CN, -OH, $-CF_3$, $-OR^6$, $-SR^6$, $-COOR^6$, the linear or branched C_1-C_{12} alkyl radicals optionally carrying an -OH, $-OR^6$ and/or -CN group, and the linear or branched C_1-C_8 alkenyl radicals;



15

20

• formula (VII)

in which:

- $\mathbf{R}^{\mathbf{5}}$, which are identical or different, have the same meanings as in formula (III),

5 - \mathbf{Y} , which are identical or different, represent X and/or \mathbb{R}^4 ,

-Z represents:

- · a direct bond,
- a C_1 - C_6 divalent alkylene radical, or a phenylene, diphenylene or phenylene-T-phenylene radical, or alternatively forms, with the two substituents R^5 and the two carbon atoms carrying these substituents, a cyclopentane or cyclohexane nucleus,
- \cdot a divalent group $-0-R^{12}-0-$, $-0-SiR^8R^9-0-SiR^8R^9-0-$, or $-0-SiR^8R^9-0-$,
- R^{12} represents a $\text{C}_2\text{-}\text{C}_8$ alkylene, $\text{C}_4\text{-}\text{C}_6$ alkenylene or xylylene radical,
- and Ar^4 has the same meaning as Ar^1 of formula (IV) in the case where n = 1.
 - family of thioxanthones of formula (VIII):

- m = 0 to 8,

10

- $\mathbf{R^{17}}$, identical or different substituent(s) on the aromatic nucleus (nuclei), represent a linear or branched C1-C12 alkyl radical, a C6-C12 cycloalkyl radical, a radical $\mathbf{Ar^1}$, a halogen atom, an -OH, -CN, -NO₂, -COOR⁶, -CHO, Ophenyl, -CF₃, -SR⁶, -Sphenyl, -SO₂phenyl, Oalkenyl, or -SiR⁶₃ group.

• family of xanthenes of formula (IX):

n = 0 to 8

family of xanthones of formula (X):

p = 0 to 8

• family of the naphthalene of formula (XI):

15 q = 0 to 8

• family of the anthracene of formula (XII):

r = 0 to 10

• family of the phenanthrene of formula (XIII):

s = 0 to 10

• family of the pyrene of formula (XIV):

5 t = 0 to 10

• family of the fluorene of formula (XV):

u = 0 to 9

• family of the fluoranthene of formula (XVI):

10

$$v = 0$$
 to 10

• family of the chrysene of formula (XVII):

w = 0 to 12

• family of the fluorene of formula (XVIII):

10

with x = 0 to 8, for example 2,7-dinitro-9-fluorenone,

• family of the chromone of formula (XIX):

with y = 0 to 6

• family of the eosin of formula (XX):

$$(R^{17})_z$$

$$Br$$

$$Br$$

$$Br$$

$$Br$$

$$Br$$

$$Br$$

with z = 0 to 5

with z = 0 to 6

• family of the erythrosin of formula (XXI):

$$(R^{17})_z$$
 $(R^{17})_z$ $(R^{17})_z$ $(R^{17})_z$

• family of the biscoumarins of formula (XXII):

with z = 0 to 5 with z = 0 to 6

$$(R^{18})_x$$
 $(R^{18})_x$

- $\mathbf{R^{18}}$, identical or different, has the same meaning as $\mathbf{R^{17}}$ or represents a group $-\mathrm{NR^6}_2$, for example 3,3'-carbonylbis(7-diethylaminocoumarin) and 3,3'-carbonylbis(7-methoxycoumarin).

Other sensitizers can be used. In particular, the photosensitizers described in the documents US 4,939,069; US 4,278,751; US 4,147,552 may be used.

In the context of the present invention, the photosensitizers have a residual absorption of UV light between 200 and 500 nm, preferably 400 to 500 nm for the preparations of dental prostheses. For dental restoration, a photosensitizer having a residual absorption of UV light above 400 nm will be preferred.

According to a preferred variant, the photosensitizers will be chosen from those of the families (IV), (VII) and (VIII). By way of examples, the following photosensitizers will be mentioned:

4,4'-dimethoxybenzoin; 2-4-diethylthioxanthone

2-ethylanthraquinone; 2-methylanthraquinone;

1,8-dihydroxyanthraquinone; dibenzoylperoxide;

2,2-dimethoxy-2-phenylacetophenone;

benzoin;

2-hydroxy-2-methylpropiophenone;

benzaldehyde;

4-(2-hydroxyethoxy)phenyl-(2-hydroxy-2-methylpropyl)-

ketone;

benzoylacetone;

2-isopropylthioxanthone;

1-chloro-4-propoxy-

thioxanthone;

4-isopropylthioxanthone; and the mixture thereof.

Various types of fillers can be used for preparing the compositions according to the invention. The fillers are chosen according to the final use of the dental composition: these affect important properties such as appearance, penetration of UV radiation, as well as the mechanical and physical properties of the material obtained after crosslinking and/or polymerization of the dental composition.

As reinforcing filler, there may be used treated or untreated pyrogenic silica fillers, amorphous silica fillers, quartz, glass or nonglassy fillers based on oxides of zirconium, barium, calcium,

15

fluorine, aluminum, titanium, zinc, borosilicates, aluminosilicates, talc, spherosil, yterbium trifluoride, fillers based on polymers in ground powder form, such as inert or functionalized methyl polymethacrylates, polyepoxides or polycarbonates.

By way of example, there may be mentioned:

- inert fillers based on methyl polymethacrylate LUXASELF from the company UGL, which can be used in the dental field and which are pigmented in pink,
- hexamethyldisilazane-treated fumed silica fillers having a specific surface area of $200\ m^2/g$,
- untreated fumed silica fillers ("aerosil" AE200 marketed by DEGUSSA).

According to an advantageous variant of the invention, the fillers and in particular the silica fillers are treated before use at 120°C with a quantity of less than 10% w/w of silicone comprising at least one unit of formula (XXIII):

$$Z - Si - (R^0)_a O_{(3-a)/2}$$

- such that Z' has the same definition as Z- a = 0,1,2 or 3
 - with at least one silicon atom.
- There may be mentioned by way of example the polymer described below with Z = epoxide and Z = trialkoxysilyl

$$(CH_3)_3Si-O = Si-O = Si-O = Si(CH_3)_3$$

$$CH_3 = Si-O = Si(CH_3)_3$$

$$CH_3 = Si(OEt)_3 = C$$

In this case for the treatment of siliconecontaining filler(s), in particular silica, with this
type of polymer, the material obtained after
crosslinking has a mechanical strength, a modulus of
elasticity and a resistance to compression which are
markedly improved.

In addition to the reinforcing fillers, pigments may be used to color the dental composition according to the invention envisaged and the ethnic groups.

For example, red pigments are used in the presence of microfibers for the dental compositions used for the preparation of dental prostheses in order to simulate the blood vessels.

Pigments based on metal oxides (iron and/or titanium and/or aluminum and/or zirconium oxides, and the like) are also used for the dental compositions used for the preparation of restoration material, in

order to obtain a crosslinked material having an ivory color.

Other additives may be incorporated into the dental compositions according to the invention. For example, biocides, stabilizers, flavoring agents, plasticizers and adherence promoters.

Among the additives which may be envisaged, there will be advantageously used crosslinkable and/or polymerizable coreagents of the organic type. These coreagents are liquid at room temperature or are hotmeltable at a temperature of less than 100°C, and each coreagent comprises at least two reactive functional groups such as oxetane-alkoxy, oxetane-hydroxyl, oxetane-alkoxysilyl, carboxyl-oxetane, oxetane-oxetane, alkenyl ether-hydroxyl, alkenyl ether-alkoxysilyl, epoxy-alkoxy, epoxy-alkoxysilyls, dioxolane-dioxolane-alcohol, and the like.

The dental compositions according to the invention may be used for numerous dental applications,

20 and in particular in the field of dental prostheses, in the field of dental restoration and in the field of temporary teeth.

The dental composition according to the invention is preferably provided in the form of a single product containing the various components ("monocomponent") which facilitates its use, in particular in the field of dental prostheses.

Optionally, the stability of this product may be provided for by organic derivatives with amine functional groups according to the teaching of the document WO 98/07798.

In the field of dental prostheses, the product in the "monocomponent" form may be deposited with the aid of a syringe directly on the plaster model or in a core. Next, it is polymerized (polymerization by possible successive layers) with the aid of a UV lamp (visible light spectrum 200-500 nm).

In general, it is possible to produce a lasting and esthetic dental prosthesis in 10 to 15 min.

It should be noted that the products obtained from the dental composition according to the invention are nonporous. Thus, after an optional polishing with the aid of a felt brush, for example, the surface of the dental prostheses obtained is smooth and bright and therefore does not require the use of varnish.

The applications in the field of dental 20 prostheses are essentially those of the joined prosthesis, which can be divided into two types:

- full prosthesis in the case of a patient with absolutely no teeth
- partial prosthesis due to the absence of several teeth, resulting either in a provisional prosthesis, or a skeleton brace.

In the field of dental restoration, the dental composition according to the invention may be used as material for filling the anterior and posterior teeth in different colors (for example "VITA" colors), which is rapid and easy to use.

The dental composition being nontoxic and polymerizable in thick layers, it is not essential to polymerize the material in successive layers. In general, a single injection of the dental composition is sufficient.

The preparations for dental prostheses and for restoration materials are carried out according to techniques which are customary in the art.

In the case of application of the dental composition as a tooth, either the tooth may be pretreated with a bonding primer or the dental composition may be prepared as a mixture with a bonding primer before its use. However, it is not essential to use a bonding primer in order to use the dental composition according to the invention.

The following examples and tests are given by way of illustration. They make it possible in particular to understand more clearly the invention and to highlight some of its advantages and to illustrate a few of its variant embodiments.

Examples and tests

The product used in the compositions of the examples are the following:

5

Product (\mathbf{B}) : this product is a mixture of siloxanes, whose viscosity is 23.5 mPa.s and in which the proportions by weight and formulae B_1 , B_2 and B_3 are given below:

10

with 89% of B_1 where a=0, 9% of B_1 where a=1; 0.2% of B_1 where a=2;

15

with 0.3% of B2 where a = 0;

and with 1.5% of B3 where a = 0 and b = 1.

5 product (P1):

Product (PS1): isopropylthioxanthone marketed under the trademark Quantacure ITX by the company RAHN.

Example 1 - Composition for dental prosthesis

The following are mixed using a three-blade stirrer:

- 100 parts of siloxane (A) stabilized with 50 ppm of Tinuvin 765;
- 1 part of photoinitiator (P1) at 75% in ethyl acetate;
 - 0.028 part of photosensitizer (PS1);
 - 150 parts of a pink-pigmented, polymethyl methacrylate-based inert filler (product LUXASELF from UGL dentaire).

The composition obtained is perfectly stable in the absence of light for several months at room temperature. This composition can be worked manually and for several hours in the presence of daylight.

A test piece 2.8 ± 0.3 mm thick is prepared in a glass dish 64 mm long (model), 10 mm wide (model) and open at the top by pouring the prepared composition ("monocomponent") into the dish.

The composition is dried by passing the dish for 1 to 2 seconds (3 m/min) under a UV lamp of 200 W/cm power corresponding to the excitation of a mixture of mercury and gallium and emitting in the UV-visible range above 400 nm.

The product obtained is unmolded by breaking the glass.

The SHORE D hardness of the two polymerized compositions is determined on each side of the item made immediately after crosslinking.

Example 1	Immediate measurement	Measurement after 10 hours
Irradiated surface:	medsar emeric	,
irradiated surface:	70	85
Bottom surface	60	85

The Shore D hardness continues to change substantially over a few hours.

2.0

The volume shrinkage is very low and excellent size stability is obtained.

The loss of mass is less than 1%.

The product may be used with or without

5 bonding primer in the presence of artificial teeth or of natural teeth.

More generally, the properties of the material obtained are in agreement with the DIN/ISO 1567 standard.

10 Example 2 - Composition for dental restoration

A composition for dental restoration is prepared by mixing:

- 200 parts of siloxane (A) stabilized with 50 ppm of Tinuvin 765;
- 1.8 parts of photoinitiator (P1) at 75% in ethyl acetate;
 - 0.0178 part of photosensitizer (PS1);
 - 52 parts of hexamethyldisilazane-treated fumed silica having a specific surface area of 200 m^2/g ;
 - 20 parts of untreated amorphous silica dried for 4 hours at 200°C before formulation.

A flowing composition is obtained which has a 25 translucent gray appearance.

The crosslinking-polymerization operation is carried out using a lamp emitting a light spot emitted

20

through a curved light tip 8 mm in diameter. The source is an Optibulb 80 W lamp (DEMETRON Optilux 500) for wavelengths of between 400 and 520 nm.

The dental composition is applied in a tooth.

5 A thickness of 5 mm is crosslinked in less than 30 seconds.

A SHORE D hardness of 50 is obtained immediately and can reach 80 to 100 within a few hours.

No loss is observed in size stability. The porosity of the crosslinked material, based on observation of a section under a microscope, is zero.

The color of the composite after crosslinking is close to ivory color.

Example 3 - Composition for dental restoration

The same concentrations of components and the same procedure as above in Example 2 are used.

However, the mixture of fillers is treated, before use, at 120°C with 5% w/w of silicone of average general formula containing less than 50 ppm of residual platinum:

$$(CH_3)_3Si-O$$
 CH_3
 CH_3

The restoration material obtained after crosslinking according to the procedure of Example 2 has a better mechanical strength, an improved modulus of elasticity and a better compression resistance.

 $\,$ A SHORE D hardness of 80 is obtained immediately.

The size stability and the porosity properties of the material are excellent.

10 Example 4 - Dental precomposition

A dental precomposition, prepared without fillers, is obtained by mixing using a magnetic stirrer bar:

- 1 part of siloxane (A) having a density of 0.997, stabilized with 50 ppm of Tinuvin 765,

- 0.01 part of photoinitiator (P1) at 10% in solution, dissolved directly in siloxane A,

- 0.00028 part of photosensitizer (PS1) contained in the photoinitiator (P1).

The crosslinking operation is carried out in a manner identical to that of Example 2. A composition 5 mm thick is crosslinked in less than 30 seconds.

The density of the crosslinked composition is measured using a brass pycnometer and its value is

1.0274. The volume shrinkage is therefore 3.05% $[=(1.0274-0.997)/0.997\times100] \ \, \text{in the absence of fillers}.$ Consequently, a dental composition of this type,

formulated with more than 50% of filler, will have a very low volume shrinkage which is less than 1.5%.

Example 5 - Dental precomposition

(a) A dental precomposition prepared without fillers is obtained by mixing using a three-blade stirrer,

- 10 parts of siloxane (A) having a density of 0.997, stabilized with 50 ppm of Tinuvin 765,

- 0.01 part of photoinitiator (P1) dissolved directly in silicone (A),

- and 62 ppm of photosensitizer (PS1) contained in the photoinitiator.

Eight grams of this dental precomposition are

then placed in an open cylindrical aluminum cup in a

manner such that the volume occupied represents a

thickness of about 6 mm.

The liquid is photocrosslinked by passing the cup under a UV lamp delivering, through a pane of glass 20 6 mm thick, UV-V (> 390 nm) = 0.6 W/cm². The UV-V (> 390 nm) dose received is 0.4 J/cm² at 10 m/min.

The degree of conversion of the epoxy

functional groups immediately after polymerization is

measured from the residual heat of reaction recorded by

25 differential calorimetry. The residual heat is 17 J/g

relative to a noncrosslinked sample which represents

237 J/g. The effective degree of conversion of the

epoxy functional groups with 0.1% photoinitiator is therefore 93%.

- (b) The same composition as above in (a) is prepared by adding 5% w/w of 3-ethyl-
- 5 3-(hydroxymethyl)oxetane during the preparation of the composition. The composition is then crosslinked in the same manner as above in (a).

The degree of conversion of the epoxy functional groups is 99.3% with a residual heat of 1.5 J/g.

(c) The same composition as above in (a) is prepared by adding 10% w/w of 3-ethyl3-(hydroxymethyl)oxetane during the preparation of the composition. The composition is then crosslinked in the

The degree of conversion of the epoxy functional groups is 99.95% with a residual heat of $0.1 \cdot J/g$.

Example 6 - Composition for dental prosthesis or dental

20 restoration material

15 same manner as above in (a).

The following are mixed using a three-blade stirrer:

- 95 parts of siloxane (A) stabilized with 50 ppm of Tinuvin 765,
- 62 ppm of photosensitizer (PS1),- 0.5 part of photoinitiator (P1) at 10% in

siloxane (A),

- 5 parts of 3-ethyl-3-(hydroxymethyl)oxetane,
 - and 120 parts of precipitated silica(ground quartz).
- A dental composition is obtained with is opaque, gray in color, nonflowing and easy to handle.

The crosslinking operation is carried out in a manner identical to that of Example 2. A composition 5 mm thick crosslinks in less than 30 seconds.

The color of the material after crosslinking is close to ivory color.

The material is suitable in particular for dental prosthesis use, in particular the stiffness is greater than 80 Mpa according to the ISO 1567 standard.

15 Example 7 - Dental composition

This composition is formulated with:

- 95 parts of silicone (B),
- 62 ppm of photosensitizer (PS1),
- 0.5 part of photoinitiator (P1) at 10% in siloxane (B),
- 5 parts of the oxetane 3-ethyl-
- 3-(hydroxymethyl)oxetane,
- and 120 parts of precipitated silica
 (ground quartz).
- 25 The crosslinking operation is carried out in a manner identical to that of Example 2.

The stiffness values found are greater than 80 Mpa according to the ISO 1567 standard.

Example 8 - Dental precomposition

A dental precomposition M1 prepared without 5 fillers is obtained by mixing using a three-blade stirrer:

> - 100 parts of a silicone composition (B) having a density of 0.997 and stabilized with 50 ppm of Tinuvin 765,

- 0.1 part of photoinitiator (P1) at 10% in solution, dissolved directly in silicone (B), - and 0.028 part of photosensitizer (PS1).

A portion of the composition obtained M1 is removed so as to crosslink it. The crosslinking 15 operation is carried out in a manner identical to that of Example 2. A composition 5 mm thick crosslinks in less than 30 seconds.

The density of the crosslinked material, measured using a brass pycnometer, is 1.0274. The 20 volume shrinkage is therefore 3.05% $[=(1.0274-0.997)/0.997 \times 100]$ in the absence of fillers. Consequently, a dental composition of this type, formulated with more than 50% of fillers, will have a very low volume shrinkage which is less than 1%.

10

Example 9 - Dental precomposition

- (a) Eight grams of **M1** of Example 8 are placed in an open cylindrical aluminum cup such that the volume occupied represents a thickness of about 6 mm.
- The photocrosslinking of **M1** and the calculation of the degree of conversion of the epoxy functional groups are carried out according to the methods described in Example 5.

The residual heat is 17 J/g relative to a

10 noncrosslinked sample which represents 237 J/g.

Consequently, the effective degree of conversion of the epoxy functional groups with 0.1% of photoinitiator is therefore 93%.

(b) The photocrosslinking and the calculation of the degree of conversion of the epoxy functional groups are also carried out for a composition M1 containing 5% w/w of a silicone oil of formula B₁ with a = 12 on average.

The degree of conversion of the epoxy

functional groups is 99% with a residual heat of

1.5 J/g.

(c) The photocrosslinking and the calculation of the degree of conversion of the epoxy functional groups are also carried out for a composition M1 containing 5% w/w of a silicone oil with a = 12 on average and of formula:

The degree of conversion of the epoxy functional groups is 99% with a residual heat of 0.1 J/g.

(d) The photocrosslinking and the calculation of the degree of conversion of the epoxy functional groups are also carried out for a composition M1 containing 5% by weight of siloxane resin of the MQM^{Epoxy} type containing 0.5% by weight of hydroxysilyl functional groups and 2% by weight of epoxy functional group (mass 43).

The degree of conversion of the epoxy functional groups is 99%.

15 Example 10 - Composition for dental prosthesis or dental restoration material

The following are mixed using a three-blade stirrer:

- 100 parts of silicone (B),
- 62 ppm of photosensitizer (PS1),
 - 0.5 part of photoinitiator (P1) at 10% in solution in silicone (B),
 - and 120 parts of precipitated silica (ground quartz).

A mixture is obtained which is opaque, gray in color, nonflowing and capable of being handled.

The crosslinking operation is carried out in a manner identical to that of Example 2. A composition 5 mm thick is crosslinked in less than 30 seconds. The color of the material after crosslinking is similar to ivory color.

The composition, in this case, is suitable in particular for dental prostheses, in particular the

10 stiffness is greater than 80 Mpa according to the

ISO 1567 standard.

Example 11 - Dental composition

The following are mixed using a three-blade stirrer:

- 95 parts of silicone (B) stabilized with 50 ppm of Tinuvin 765,
 - 62 ppm of photosensitizer (PS1),
 - 0.5 part of photoinitiator (P1) at 10% in silicone (B),
- 20 5 parts of polydimethylsiloxane **B1** where a
 - = 12 on average,
 - and 120 parts of precipitated silica (ground quartz).

The crosslinking operation is carried out in 25 a manner identical to that of Example 2. The measured stiffness values are greater than 80 Mpa according to the ISO 1567 standard.

CLAIMS

- 1. A dental composition comprising:
- (1) at least one crosslinkable and/or polymerizable silicone oligomer or polymer which is liquid at room temperature or which is heat-meltable at a temperature of less than 100°C, and which comprises:
 - at least one unit of formula (FS):

$$Z - Si - (R^0) O_{(3-a)/2}$$

10

15

20

25

in which:

- -a = 0, 1 or 2,
- R^0 , identical or different, represents an alkyl, cycloalkyl, aryl, vinyl, hydrogeno or alkoxy radical, preferably a C_1 - C_6 lower alkyl,
- Z, identical or different, is an organic substituent comprising at least one reactive epoxy, and/or alkenyl ether and/or oxetane and/or dioxolane and/or carbonate functional group,
- and at least two silicon atoms,
- (2) at least one aromatic hydrocarbon photosensitizer with one or more aromatic nuclei which are substituted or not, having a residual light absorption of between 200 and 500 nm,

10

15

20

- (3) at least one dental filler present in a proportion of at least 10% by weight relative to the total weight of the composition,
- (4) and an effective quantity of at least one borate-type photoinitiator, chosen from those of formula:

 Δ in which the cationic entity of the borate is selected from:

(*) the onium salts of formula (I): $[(R^{1})_{n}-A-(R^{2})_{m}]^{+}$ (I)

in which formula:

- A represents an element of groups 15 to 17 such as for example: I, S, Se, P or N,
- R^1 represents a carbocyclic or heterocyclic C_6 - C_{20} aryl radical, it being possible for said heterocyclic radical to contain, as heteroelements, nitrogen or sulfur,
- R^2 represents R^1 or a linear or branched C_1 - C_{30} alkyl or alkenyl radical; said radicals R^1 and R^2 being optionally substituted with a C_1 - C_{25} alkoxy, C_1 - C_{25} alkyl, nitro, chloro, bromo, cyano, carboxyl, ester or mercapto group,
 - n is an integer ranging from 1 to v+1, v being the valency of the element A,
- m is an integer ranging from 0 to v 1 with n + m = v + 1,
 - (**) the organometallic salts of formula (III):

20

25

$(L^1L^2L^3M)^{+q}$

in which formula:

- M represents a group 4 to 10 metal, in particular iron, manganese, chromium or cobalt,
- L¹ represents 1 ligand bound to the metal M by π by electrons, which ligand is chosen from the ligands η^3 -alkyl, η^5 -cyclopendadienyl and η^7 -cycloheptratrienyl and the η^6 -aromatic compounds chosen from the optionally substituted η^6 -benzene ligands and the compounds having from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 3 to 8 π electrons;
 - L² represents a ligand bound to the metal M by π electrons, which ligand is chosen from the ligands η^7 -cycloheptratrienyl and the η^6 -aromatic compounds chosen from the optionally substituted ligands η^6 -benzene and the compounds having from 2 to 4 condensed rings, each ring being capable of contributing to the valency layer of the metal M by 6 or 7 π electrons;
 - L^3 represents from 0 to 3 ligands, which are identical or different, linked to the metal M by σ electrons, which ligand(s) is (are) chosen from CO and NO_2^+ ; the total electron charge q of the complex to which L^1 , L^2 and L^3 contribute and the

ionic charge of the metal M being positive and equal to 1 or 2;

 Δ the anionic borate entity of which has the formula $[BX_aR_b]^-$ in which:

- 5 a and b are integers ranging, for a, from 0 to 3 and, for b, from 1 to 4 with a + b = 4,
 - the symbols X represent:
 - * a halogen atom (chlorine, fluorine) with a=0 to 3,
- 10 * an OH functional group with a = 0 to 2,
 - the symbols R are identical or different and represent:
 - \gt a phenyl radical substituted with at least one electron-attracting group such as for example OCF₃, CF₃, NO₂, CN, and/or with at least 2 halogen atoms (fluorine most particularly), this being when the cationic entity is an onium of an element of groups 15 to 17,
- P a phenyl radical substituted with at least one
 20 element or one electron-attracting group, in
 particular a halogen atom (fluorine most
 particularly), CF3, OCF3, NO2, CN, this being when
 the cationic entity is an organometallic complex
 of an element of groups 4 to 10,
- 25 > an aryl radical containing at least two aromatic nuclei such as for example biphenyl, naphthyl, optionally substituted with at least one electron-

attracting group or element, in particular a halogen atom, including fluorine in particular, OCF_3 , CF_3 , NO_2 , CN, regardless of the cationic entity.

- 2. The composition as claimed in claim 1, characterized in that Z is an organic substituent Z1 comprising at least one reactive epoxy, and/or dioxolane functional group, and preferably at least one reactive epoxy functional group.
- 3. The composition as claimed in claim 2, characterized in that the oligomer or polymer (1) comprises in addition other reactive functional groups Z such as the reactive alkenyl ether, oxetane and/or carbonate functional groups Z2.
- 4. The composition as claimed in any one of the preceding claims, characterized in that the reactive functional group(s) of Z1 are chosen from the following radicals:

The composition as claimed in any one of 5. the preceding claims, characterized in that the photoinitiator is chosen from the group consisting of: $[(C_8H_{17})-O-\Phi-I-\Phi)]^+$, $[B(C_6F_5)_4]^ [(\Phi)_2 \ I]^+$, $[B(C_6F_5)_4]^ [C_{12}H_{25}-\Phi-I-\Phi]^+$, $[B(C_6F_5)_4]^ [(C_8H_17-O-\Phi)_2I]^+$, $[B(C_6F_5)_4]^ [(C_8H_17)-O-\Phi-I-\Phi)]^+$, $[B(C_6F_5)_4]^ [(\Phi)_3S]^+$, $[B(C_6F_5)_4]^ [(\Phi)_2S-\Phi-O-C_8H_{17}]^+$, $[B(C_6H_4CF_3)_4]^-[(C_{12}H_{25}-\Phi)_2I]^+$, $[B(C_6F_5)_4]^ [(\Phi)_3 S]^+, [B(C_6F_4OCF_3)_4]^ [(\Phi-CH_3)_2I]^+$, $[B(C_6F_5)_4]^ [(\Phi-CH_3)_2 I]^+, [B(C_6F_4OCF_3)_4]^ [CH_3-\Phi-I-\Phi-CH(CH_3)_2]^+, [B(C_6F_5)_4]^ (\eta^5$ -cyclopentadienyl) $(\eta^6$ -toluene) Fe $^+$, $[B(C_6F_5)_4]^-$ 5 $(\eta^5\text{-cyclopentadienyl})(\eta^6\text{-methyl-1-naphthalene})$ Fe $^+$, $[B(C_6F_5)_4]^{-1}$ $(\eta^5$ -cyclopentadienyl) $(\eta^6$ -cumene) Fe⁺, [B(C₆F₅)₄]⁻ and the mixture thereof. 10 The composition as claimed in any one of

the preceding claims, characterized in that the

photosensitizer is chosen from the group consisting of:

4,4'-dimethoxybenzoin; 2-4-diethylthioxanthone

2-ethylanthraquinone; 2-methylanthraquinone;

1,8-dihydroxyanthraquinone; dibenzoylperoxide;

2,2-dimethoxy-2-phenylacetophenone;

benzoin;

2-hydroxy-2-methylpropiophenone;

benzaldehyde;

4-(2-hydroxyethoxy)phenyl-(2-hydroxy-2-methylpropyl)-ketone;

benzoylacetone;

2-isopropylthioxanthone;

1-chloro-4-propoxy-

thioxanthone;

4-isopropylthioxanthone;

and the mixture thereof.

7. The dental composition as claimed in any one of the preceding claims, characterized in that the silicone oligomer and/or polymer (1) consists of at least one silicone having the following average formula:

CHCH CH2

CH2CH2CH2

CH3 Si-CH3

CH3 Si-CH3

CH2CH2CH2

CH2CH2CH2

- 8. The use of a dental composition as
 5 claimed in any one of the preceding claims for the
 production of dental prostheses.
 - 9. The use of a dental composition as claimed in any one of claims 1 to 7, for dental restoration.
- 10. A dental prosthesis which can be obtained from a composition as claimed in any one of claims 1 to 7.
- 11. A dental restoration material which can be obtained from a composition as claimed in any one of 15 claims 1 to 7.

DENTAL COMPOSITION BASED ON SILICONE CROSSLINKABLE BY CATION PROCESS

5

Abstract of the Disclosure

The invention concerns dental compositions. Said composition comprises (1) a silicone crosslinkable

10 and/or polymerizable by cation process; (2) an efficient amount of at least an initiator such as onium borate; (3) at least a photosensitizer; and (4) a dental filler present in the composition in a proportion of at least 10 wt. % relative to the

15 composition total weight. Said dental compositions are useful for making dental prostheses or for dental restoration.

20 C/Data/RN98131.Abst

Express	Mail	l ahel	#FI	610	2390	1221	2111	
	IVICIL	Lavei	TT L L.		JUJ3	, coco		

		AND POWER OF ATTORNEY	RN98131	DOCKET NO
(include Reference to PCT Inte	rnational Applications) PC 1/	F K37/U4343	KN98131	
As a below named inventor, 1	hereby declare that:	•		
My residence, post office addi	ress and citizenship are as sta	ted below next to my name.		
I believe I am the original, first are listed below) of the subjec	and sole inventor (if only one that the matter which is claimed and	name is listed below) or an origin for which a patent is sought on th	nal, first and joint ne invention entitle	inventor (if plural names ed:
DENTAL COMPOS PROCESS	ITION BASED ON S	BILICONE CROSSLIN	KABLE BY	CATION
the specification of which (che	eck only one item below):			
is attached here	to.			
-	ted States application			
Serial No.				
on				
and was amende	•			
		(if applicable)		
_	T international application			
Number PCT/	FR99/02345		 _	
on <u>Octobe</u>	1st, 1999			
and amended u	nder PCT ARTICLE 19			
on	(if applicable).			
I hereby state that I have revi by any amendment referred t		tents of the above-identified spec	cification, including	g the claims, as amended
I acknowledge the duty to dis Code of Federal Regulations.	close information which is mail	terial to the patentability of this a	oplication in accor	rdance with Title 37,
I hereby claim foreign priority	henefits under Title 35. Unite	d States Code, §119 of any foreig	nç	
application(s) for patent or in	ventor's certificate or of any Policeted below and have also ide	CT international application(s) de entified below any foreign applica	signating at least ition(s) for patent	or inventors certificate or
any PCT international applica	ation(s) designating at least on	e country other than the United S tion(s) of which priority is claime	States of America	filed by me on the same
Subject matter naving a ming	date beidle frat or the abbilde	dion(s) of which phoney to country	- .	
PRIOR FOREIGN/PCT APP	LICATION(S) AND ANY PRICE	PRITY CLAIMS UNDER 35 U.S.C	c. 119:	
COUNTRY	APPLICATION NUMBER	DATE OF FILING	PRIOR	ITY CLAIMED
PCT indicate PCT		(day month year)	INDER 35 USC 1	19
FRANCE	98/12375	02 October, 1998	∠ YES	NO
			YES	NO
			YES	□ NO
			YES	□ NO
	DACE 4 of 2	LIS DEPARTMENT OF C	YES	□ NO

OMBI Contir	NED DECLARATI nued) (include Refer	ON FOR PA	TENT APPLICATION A International Applications	AND POWER OF ATTO S) PCT/FR99/0234	DRNEY 15		RN98131
oplica f this a tates 1.56(a pplica	tion(s) designating application is not di Code, §112, I acknument which occurred button:	the United S isclosed in the lowledge the between the f	5, United States Code, tates of America that is at those prior applicatio duty to disclose materia iling date of the prior ap	lare listed below and, on(s) in the manner pro al information as define oplication(s) and the na	insofar as the ovided by the feed in Title 37, ational or PCT	subject matter of first paragraph o Code of Federa international fili	or each of the claims if Title 35, United Il Regulations. ing date of this
RIOR		NS OR PCT	INTERNATIONAL APP	PLICATIONS DESIGN	ATING THE U	J.S. FOR BENE	FIT UNDER 35
<u>.3.0.</u>	120	U.S. APPLIC	ATIONS		TUS (CHECK		Linupourn
U.S	S. APPLICATION NUMBER		U.S. FILING DATE	PATENTE	ED	PENDING	ABANDONED
PCT	APPLICATIONS I	DESIGNATIN	IG THE U.S.				
		CT FILING DATE	US SERIAL NUMBE ASSIGNED (if any				
b							
JEAN	Correspondence	to: IET		tion under 37 CFR	Direct Tel (name and telep JEAN-LC	ephone Cails to hone number) OUIS SEUGNE	
JEAN INTE RHO 259 F	Correspondence	to: IET OPERTY DE INS ROAD	- Limited Recognit		Direct Tel	ephone Calls to hone number) DUIS SEUGNE 0-4180	iT
JEAN INTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR	to: IET OPERTY DE INS ROAD Y, NJ 0851	F - Limited Recognit	FIRST GIVEN Jean-Marc	Direct Tel (name and telep JEAN-LC (609) 86	ephone Calls to hore number) OUIS SEUGNE 0-4180	IAME
JEAN INTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE &	to: IET OPERTY DE INS ROAD Y, NJ 0851	EPT.	FIRST GIVEN	Direct Tel (name and telep JEAN-LC (609) 86	ephone Calls to home number) OUIS SEUGNE 0-4180 SECOND GIVEN N	IAME IZENSHIP
NTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR	to: IET OPERTY DE INS ROAD Y, NJ 0851 FAMILY NAM FRANCI CITY MEYZIE POST OFFICE	EPT. 12-7500 1EE ES	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL	Direct Tel (name and telep JEAN-LC (609) 86	ephone Calls to home number) OUIS SEUGNE 0-4180 SECOND GIVEN N	IAME IZENSHIP DE/COUNTRY
NTERHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR	to: IET OPERTY DE INS ROAD Y, NJ 0851 FAMILY NAM FRANCI CITY MEYZIE POST OFFICE	EPT. 12-7500 IE ES EADDRESS	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FRANCE	Direct Tel (name and telep JEAN-LC (609) 86	SECOND GIVEN N COUNTRY OF CIT FRANCE STATE & ZIP COD	IAME IZENSHIP DE/COUNTRY
NTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP	to: IET OPERTY DE INS ROAD Y, NJ 0851 FAMILY NAM FRANCI CITY MEYZIE POST OFFICE	EPT. 12-7500 IE ES EADDRESS	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FRANCE	Direct Tel (name and telep JEAN-LC (609) 86	SECOND GIVEN N COUNTRY OF CIT FRANCE STATE & ZIP COD	IAME IZENSHIP DE/COUNTRY
NTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE &	to: IET OPERTY DE INS ROAD Y, NJ 0851 FAMILY NAM FRANCI CITY MEYZIE POST OFFIC 1, rue de	EPT. 12-7500 IE ES IU EADDRESS Flandres	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FRANCE CITY MEYZIEU	Direct Tel (name and telep JEAN-LC (609) 86	SECOND GIVEN N COUNTRY OF CIT FRANCE STATE & ZIP COD F-69330, F	IAME TIZENSHIP DE/COUNTRY RANCE
JEAN NTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR	FAMILY NAME FAMILY	EPT. 12-7500 IE ES IU EADDRESS Flandres	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FRANCE CITY MEYZIEU FIRST GIVEN	Direct Tel (name and telept JEAN-LC (609) 86	SECOND GIVEN I	IAME IZENSHIP DE/COUNTRY RANCE
JEAN NTE RHO 259 F CN 7	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP RESIDENCE & CITIZENSHIP	FAMILY NAME OF THE POST OF THE	EPT. 12-7500 IE ES Flandres	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FIRST GIVEN FIRST GIVEN STATE OR FOREIGN CO	Direct Tel (name and telept JEAN-LC (609) 86	SECOND GIVEN N SECOND GIVEN N COUNTRY OF CIT FRANCE STATE & ZIP COD F-69330, FI SECOND GIVEN N COUNTRY OF CIT	IAME IZENSHIP DE/COUNTRY RANCE NAME
JEAN NTE RHO 259 F 201 /	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE	FAMILY NAME OF THE POST OF FRENCE OF THE POST OF FRENCE OF THE OFFICE OF THE OFFI	EPT. 2-7500 EE ES Flandres CE ADDRESS	FIRST GIVEN Jean-Marc STATE OR FOREIGN COL FRANCE CITY MEYZIEU FIRST GIVEN STATE OR FOREIGN COL CITY	Direct Tel (name and telep) JEAN-LC (609) 86	SECOND GIVEN IN SECOND GIVEN	ET LAME TIZENSHIP DE/COUNTRY RANCE NAME TIZENSHIP DE/COUNTRY
202 203	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS eby declare that all believed to be true:	FAMILY NAME OF STATE	EPT. 2-7500 EE ES Flandres THE CE ADDRESS made herein of my own hat these statements we isonment or both, under the company of the compa	FIRST GIVEN Jean-Marc STATE OR FOREIGN COLFRANCE CITY MEYZIEU FIRST GIVEN STATE OR FOREIGN COLTY I knowledge are true at the knowledge are true at the section 1001 of Title	Direct Tel (name and telep) JEAN-LC (609) 86 UNTRY	SECOND GIVEN I SECOND GIVEN I COUNTRY OF CIT FRANCE STATE & ZIP COD	IAME TIZENSHIP DE/COUNTRY RANCE NAME TIZENSHIP DE/COUNTRY In information and be ments and the like so
JEAN INTE RHO 259 F CN 7 201 202	Correspondence I-LOUIS SEUGN LLECTUAL PRO DIA INC. PROSPECT PLA 500, CRANBUR FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS FULL NAME OF INVENTOR RESIDENCE & CITIZENSHIP POST OFFICE ADDRESS eby declare that all believed to be true:	FAMILY NAME OF STATE	EPT. 2-7500 EE ES Flandres CE ADDRESS	FIRST GIVEN Jean-Marc STATE OR FOREIGN COLFRANCE CITY MEYZIEU FIRST GIVEN STATE OR FOREIGN COLTY I knowledge are true at the knowledge are true at the section 1001 of Title	Direct Tel (name and telep) JEAN-LC (609) 86 UNTRY	SECOND GIVEN I SECOND GIVEN I COUNTRY OF CIT FRANCE STATE & ZIP COD	IAME TIZENSHIP DE/COUNTRY RANCE NAME TIZENSHIP DE/COUNTRY In information and ments and the like

DATE

DATE

May 12th

2001